Ultrahigh-throughput screening in drop-based microfluidics for directed evolution

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The explosive growth in our knowledge of genomes, proteomes, and metabolomes is driving ever-increasing fundamental understanding of the biochemistry of life, enabling qualitatively new studies of complex biological systems and their evolution. This knowledge also drives modern biotechnologies, such as molecular engineering and synthetic biology, which have enormous potential to address urgent problems, including developing potent new drugs and providing environmentally friendly energy. Many of these studies, however, are ultimately limited by their need for even-higher-throughput measurements of biochemical reactions. We present a general ultrahigh-throughput screening platform using drop-based microfluidics that overcomes these limitations and revolutionizes both the scale and speed of screening. We use aqueous drops dispersed in oil as picoliter-volume reaction vessels and screen them at rates of thousands per second. To demonstrate its power, we apply the system to directed evolution, identifying new mutants of the enzyme horseradish peroxidase exhibiting catalytic rates more than 10 times faster than their parent, which is already a very efficient enzyme. We exploit the ultrahigh throughput to use an initial purifying selection that removes inactive mutants; we identify ∼100 variants comparable in activity to the parent from an initial population of ∼10<sup>8</sup>. After a second generation of mutagenesis and high-stringency screening, we identify several significantly improved mutants, some approaching diffusivity-limited efficiency. In total, we screen ∼10<sup>9</sup> individual enzyme reactions in only 10 h, using <150 μL of total reagent volume; compared to state-of-the-art robotic screening systems, we perform the entire assay with a 1,000-fold increase in speed and a 1-million-fold reduction in cost.

protein engineering | compartmentalization | emulsion | horseradish peroxidase

Much of the significant progress in our ever-advancing knowledge of biology is driven by our ability to perform very large numbers of reactions and to screen large amounts of data (1). This is certainly the case for the explosion in our knowledge of the genome and of proteomes and metabolomes. It is also the case for many important emerging biotechnologies, such as molecular engineering and synthetic biology. A particularly powerful tool for these technologies is directed evolution (2–4), which is a means of producing proteins and nucleic acids with tailor-made activities, and which has led to new classes of drugs (2), as well as improved enzymes (3) and strains of microorganisms for industrial applications (4). Directed evolution is also a powerful method to study the fundamentals of evolution itself (5). Evolution is practiced in the laboratory by artificially inducing mutations to create populations of new variants. These populations are subjected to a controlled pressure allowing selection of those rare variants which exhibit improved functionality. Evolutionary information is encoded in the genotype, whereas selection acts on the phenotype; this requires the genotype and phenotype to be linked. However, to be truly effective, directed evolution requires screening or selection of very large populations, or libraries, using conditions that closely match the desired functionality. Numerous technologies can link genotype and phenotype for the selection of high-affinity binding; affinity purification can select libraries as large as 10<sup>10</sup> nucleic acid variants by the systematic removal of ligands by exponential enrichment (SELEX) (6) or 10<sup>15</sup> protein variants by phage display (7). By contrast, it is much more challenging to select for other functionalities, such as catalysis (8). Cells also can be used to link genotype and phenotype; both growth-based selections and screening cells by FACS can potentially access libraries larger than 10<sup>8</sup>, but the range of reactions is very limited, either to survival or a cell-bound product, respectively (3, 9, 10). Genotype and phenotype can also be linked by compartmentalizing individual elements of the library in the aqueous drops of a water-in-oil emulsion (9). Bulk emulsions (3, 9) allow screening of libraries as large as 10<sup>11</sup>, but the lack of control of reaction volumes, timing, and generality ultimately limit their flexibility. To achieve full control and flexibility, current state-of-the-art high-throughput screening methods employ microtiter plates and sophisticated robots to process up to 100,000 assays a day, or ~1/s. This is a thousand times more than were processed in an entire week in 1990 (11). However, this technology is approaching its physical limit; below the 1-μL-volumes of 1,536-well plates, evaporation and capillary forces become prohibitive (11). New developments exploiting microfluidic lab-on-a-chip technology are significantly improving screening capabilities, decreasing the reaction volume by roughly 1,000-fold while increasing the speed by ~10-fold (12).

Nevertheless, these screening methods still limit the applicability of directed evolution and many of the other emerging biotechnologies (13). Any further significant extension of screening capabilities will require revolutionary improvements, and these have, as yet, not been forthcoming.

In this paper, we report the development of an ultrahigh-throughput screening platform using drop-based microfluidic devices that provides the revolutionary improvements required to significantly advance our screening capabilities. We demonstrate...
the power of this platform by using it for directed evolution to
discover variants of the enzyme horseradish peroxidase (HRP).
We use picoliter-volume aqueous drops dispersed in an inert
oil as reaction vessels; most contain no more than a single yeast
cell that displays a variant of the enzyme on its surface. We sort
these drops at rates of thousands per second, allowing us to
screen a library of $10^9$ in about 10 h, using a total reagent volume
of $<150 \mu$L. This enables us to discover variants of HRP that are
more than 10-fold faster than the parent and that approach the
diffusion-limited rate. Compared to state-of-the-art robotic
screening, this is 1,000-fold faster and uses 10-million-fold less
volume of reagent, representing a cost savings estimated to be
about 10-million-fold.

**Results and Discussion**

Our ultrahigh-throughput-screening platform consists of two
polydimethylsiloxane (PDMS) microfluidic devices (14) con-
ected by an incubation line (Fig. 1). In the first device, we coflow
a suspension of yeast cells displaying enzyme variants on their
surface (10) with a second stream containing a fluorogenic sub-
strate for the enzyme. We use a long and narrow channel to pro-
duce drops with $>64\%$ aqueous volume fraction. The reaction is
initiated upon formation of these drops as the two streams mix.
The 6-pL, 23-μm-diameter, aqueous drops are dispersed in fluo-carbon oil and stabilized by a biocompatible surfactant (15).
The cells are encapsulated individually in drops (16), allowing
us to separately probe each member of the library; moreover,
the uniform drop size ensures accurate comparison of the fluo-
rescent intensities, and thus the enzymatic reaction rates. The
drops leave the first device and enter the incubation line; at a
high volume fraction, the drops are packed such that they flow
as a solid plug through the delay line. This enables us to use a
continuous process with low time dispersion during the
incubation.

The drops then flow into the second microfluidic device, a
high-speed drop sorter. We dielectrophoretically sort (17, 18)
the drops based on a preset fluorescence intensity threshold,
thereby selecting the most active enzyme variants. In the sorting
device (Fig. 1E), the gradual constriction just before the addition
of oil to the sorting device allows the drops to pack efficiently into
a single-file line, thus allowing periodic reinjection and even
spacing of the drops as they enter the sorting junction. The large
radius of curvature of the sorting junction serves to reduce shear
on the drops as they enter the keep or waste branches. This shear
ultimately limits the sorting rate in this design, as higher flow
rates will split drops at the junction. Finally, the series of arc-
shaped channels running between the waste and keep channels
serve as pressure shunts to equalize the pressure immediately
after the sorting junction. This passive method is simple, reliable,
and decouples the critically balanced flow at the sorting junction
from any disturbances downstream in the device, such as fluctu-
ating numbers of drops in each branch and the removal and in-
sertion of tubes for collection. These design considerations lead
to a robust platform that is able to sort at rates of $>2$ kHz.

The detection limit of this system is $\sim 1$ nm Amplex Ultrared
(AUR) ($\sim 3,500$ molecules in 6 pL). There are $\sim 10^7$ enzyme
molecules displayed per cell, and therefore we can detect $<1$ turn-
over per enzyme. Drops are stable to coalescence and cells are
viable for days, so it is possible to detect enzymes with rates of
less than one turnover per day. Therefore, the detection system
does not limit us from identifying the very poor catalysts expected
in early generations in an experiment to evolve new function.
Here we demonstrate sensitive detection of fluorescent products;
it is possible to use coupled assays to adapt many important en-
zyme assays to yield fluorescent products. However, with minimal
changes to the optical system, we can adapt the design to access
other modes of detection, such as luminescence and absorbance.

Improving the catalytic power of horseradish peroxidase is
challenging because it is already a very efficient enzyme (19),
especially for the substrate we use, AUR (20). We therefore
adopt a strategy that first uses a purifying selection (21) on a large
library of variants (10) (SI Text), $\sim 10^7$; the applied pressure
selects only those variants with activities similar to or better than
that of the wild-type enzyme, purging the library of inactive

**Fig. 1.** Modules of the ultrahigh-throughput microfluidic screening platform. (A) A low-magnification image of the entire drop-making device. (B) A suspension of yeast cells displaying the HRP on their surface (aq1) is combined with a second aqueous stream containing the fluorogenic substrate AUR (aq2). The yeast are at a concentration of 1 × 10^8 cells per milliliter, which gives an average of 0.3 cells per 6-pL drop after being diluted by half by the substrate stream. The aqueous drops are formed at a flow-focusing junction (33) in a fluorocarbon oil at a rate of 2 kHz, and the number of cells per drop follows a Poisson distribution; $\sim 22\%$ have a single cell (16) (SI Text). (C) The drops flow out of this device into a tube that acts as an incubation line where they incubate for $\sim 5$ min. We use a fluorosurfactant (15) to prevent coalescence and to give the drops a biocompatible interface. (D) A single layer of drops after incubation showing the fluorescence developing from the active HRP displayed on the surface of the cells. (E) From the delay line, the drops flow as a solid plug to a junction where oil is added to separate the drops. To visually demonstrate the sorting process, we sorted an emulsion containing light and dark drops; the light drops contain 1 mM fluorescein, and the dark contain 1% bromophenol blue. Fluorescence levels are detected as the drops pass a laser focused on the channel at the gap between two electrodes. When sorting is on, the light drops, which are brighter than the threshold level, are sorted by dielectrophoresis (17, 18) into the bottom channel (SI Text). (Scale bar, 80 μm.)
mutants (Fig. 2). The rationale of this low-stringency approach is that the resulting variants of the first generation may contain potentiating mutations, which do not show improved phenotype on their own, but instead may poise the enzyme for improved function upon acquiring other, functionally adaptive mutations (22, 23). Mutations that improve catalysis may be deleterious to other functions, such as structural stability and, to be effective, require the preexistence of stabilizing mutations to compensate (21).

To perform each round of screening, we induce the yeast to express the surface-displayed HRP, encapsulate the cells in drops, and sort drops at 2,000/s for up to 3 h, screening as many as 2 × 10^7 cells. Before round one of the screen, we create two libraries each containing >10^7 mutants of the HRP gene. Because mutations both near the active site and distant from it are known to affect enzyme function (24), we create two types of libraries: We use random mutagenesis by error-prone PCR (epPCR) to target residues throughout the protein. Mutagenesis by epPCR will tend to bias mutations distant from the active site (24), and thus we create a second library using saturation mutagenesis to target specific residues at the entrance to the active site where the substrate binds (Protein Data Bank code 1HCH) (25). No additional mutations are added between rounds. We assay the enrichment of the pool of selected yeast after each round (SI Text). Although there is very little detectable activity in the unselected libraries, by the fourth round, the selected population exhibits higher activity than that of the parent by a factor of ~2 (Fig. 3).

To identify the amino acid changes in the selected variants, we sequence a total of 82 individual mutants and find 50 unique sequences. All of them are full-length genes with no internal stop codons. There are between one and five amino acid substitutions per mutant gene, and all encode active enzymes with catalytic rates similar to that of the parent (Fig. 3 and SI Text). From the distributions of the numbers of identical sequences (26), we estimate there are a total of ~200 unique active sequences in the starting library of 2 × 10^6 (Fig. 3), thus only one in 10^6 of the original mutants is active (17).

The results of this purifying selection can be used to find mutants with enhanced catalytic activity. To accomplish this, we further mutate the 18 highest-activity clones from the first generation to create a new library again containing ~10^7 variants. We screen as before, but sort the brightest 5–10% of the drops, which contain the enzymes with the highest catalytic activity. The activity of the selected population reaches a plateau of ~8 times that of the parent by the fifth round of screening (Fig. 3A). We then plate the cells, measure the activity of 184 mutants, and sequence the top 50%. Of these 92 sequences, only 31 are unique (SI Text); because many sequences are identified multiple times, this indicates that the population is beginning to converge on a smaller set of highly active sequences. In contrast to the variants selected in the first generation of evolution, the second-generation mutants are all at least five times faster than the wild type, with the best being 12 times faster (Fig. 3A and SI Text). An analysis of the turnover number, k_cat, and the Michaelis constant, K_M, shows that all of the top 13 mutants from the second generation have significantly increased catalytic efficiency (k_cat/K_M). All are more than twice as efficient as the wild type, and the best mutant is nearly seven times more efficient, with k_cat/K_M ~ 2.5 × 10^5 M^-1 s^-1; this is approaching the maximum rate allowed by diffusion-limited encounters of ~10^8 M^-1 s^-1 (SI Text). Even though the screen was performed at a substrate concentration of 100 μM, near the wild-type K_M, we find that the K_M increases for all mutants; however, this is compensated for by even larger increases in k_cat, of up to 11-fold, yielding the higher catalytic efficiencies observed.

Fourteen out of 30 amino acid substitutions found in mutants from the neutral selection are inherited in mutants selected from...
the second-generation libraries. In addition, there are 27 new substitutions in the second-generation sequences, giving a total of 41 substitutions. Every second-generation mutant inherits at least one substitution from the first generation, although as expected, none of the first-generation sequences occur exactly in the second generation, because a much higher threshold is used for sorting. Of the 31 unique sequences, 23 contain at least one new substitution, while the remaining eight contain only recombinations of the first-generation substitutions. Interestingly, some substitutions that give only moderate increases in catalytic rates in first-generation mutants are also found in many of the fastest second-generation mutants. Seven of the 14 inherited substitutions cluster around sites of known structural importance, such as the two structural calcium ions and Phe221, which is known to be involved in heme organization (19) (Fig. 3). The residues most frequently found mutated tend to cluster around the two structural pink calcium ions and at the opening to the active site in front of the brown heme group. Yellow residues are inherited between rounds. Note the clusters of inherited residues around the two structural pink calcium ions and at the opening to the active site; the site of substrate binding.

We quantified the advantages of the drop-based microfluidic platform by comparing requirements for the full screen to a conservative estimate for those of a robot (Table 1). A reasonable estimate for the throughput of the robot gives a total time for the screen of nearly 2 years; by comparison, the microfluidic device requires only 5 h for the full screen. This is well over a 1,000-fold reduction. Similarly, using a reaction volume of 100 µL per assay with the robot, the total volume of reagent is ∼10^8 L; by comparison, the microfluidic device uses only 150 µL of reagents. This is more than a 10-million-fold reduction. Including all supplies and amortization, the total cost for screen with the robot would be ∼1.5 million; by comparison, the cost for the microfluidic screen is under $4. This is a 4-million-fold reduction.

Table 1. Comparison of time and costs* for the complete screen using traditional methods and in microfluidic emulsions

<table>
<thead>
<tr>
<th></th>
<th>Robot</th>
<th>Microfluidic drops</th>
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<tbody>
<tr>
<td>Total reactions</td>
<td>5 × 10^7</td>
<td>5 × 10^7</td>
</tr>
<tr>
<td>Reaction volume</td>
<td>100 µL</td>
<td>6 µL</td>
</tr>
<tr>
<td>Total volume</td>
<td>5,000 L</td>
<td>150 µL</td>
</tr>
<tr>
<td>Reactions/day</td>
<td>73,000</td>
<td>1 × 10^6</td>
</tr>
<tr>
<td>Total time</td>
<td>~2 years</td>
<td>7 h</td>
</tr>
<tr>
<td>Number of plates/devices</td>
<td>260,000</td>
<td>2</td>
</tr>
<tr>
<td>Cost of plates/devices</td>
<td>$520,000</td>
<td>$1.00</td>
</tr>
<tr>
<td>Cost of tips</td>
<td>$10 million</td>
<td>$0.30</td>
</tr>
<tr>
<td>Amortized cost of instruments</td>
<td>$280,000</td>
<td>$1.70</td>
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<tr>
<td>Substrate</td>
<td>$4.75 million</td>
<td>$0.25</td>
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<tr>
<td>Total cost</td>
<td>$15.81 million</td>
<td>$2.50</td>
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*Details in SI Text
The ability to screen libraries of $>10^7$ in just a few hours at a cost of only a few dollars will be of enormous benefit for directed evolution. There has already been some success screening small libraries that yield only modest improvements, and then performing repeated rounds of mutation and screening (27). However, when selecting for the binding activity of proteins, a clear relationship between library size and the affinity of the selected proteins is observed experimentally (7): Using antibody V-genes from nonimmunized donors, small phage-antibody libraries of $<10^8$ genes yield only antibodies with $K_d \sim 10^{-10}$ affinities, whereas larger libraries of $>10^6$ yield $K_d \sim 10^{-9}$. Similar improvements in the catalytic efficiency of enzymes should be possible with the use of larger libraries hitherto impossible using traditional robotic screening systems.

The drop-based microfluidic platform described here represents a unique class of screening system. When used with cells, the system operates as a drop-based FACS in that it interrogates individual cells and sorts them based on the results. However, unlike a traditional FACS, the cells remain encapsulated in drops and the entire reaction vessel is assayed and sorted. Prior to sorting, drops can be fused (28) to add additional reagents or even other cells, further increasing the versatility. In addition, this methodology is not limited to reactions with cells (15) or even to biological reactions. In this sense, drop-based microfluidics has many of the merits of both screening using microtiter plates and FACS: The assays are compartmentalized but the compartments themselves are screened and sorted at high speed based on fluorescence. This has the potential to significantly enhance the utility and applicability of high-throughput screening, not only for directed evolution, but also in many other areas of biology and chemistry where large parameter spaces need to be explored quickly and affordably. Moreover, the improvements in the ultra-high-throughput capabilities of the drop-based microfluidics are somewhat analogous to those of microprocessors, where transistor density has increased by $\sim 100,000$-fold in the past few decades (29). These improvements have led to completely new and unexpected applications for microprocessors; so too should the improvements in screening technology lead to unexpected new applications.

Materials and Methods

Construction of HRP Libraries. For the first generation of evolution, we use two previously created libraries (10, 30). In the first, the HRP gene is randomly mutated using epPCR at a rate of one to three mutations per gene. In the second, we combinatorially mutate, using saturation mutagenesis, five residues near the mouth of the active site where the substrate binds. We target Phe68, Gln69, Ser73, and Ala74 in one reaction and positions Phe137, Leu138, Ala140, Phe142, and Phe143 in a second reaction. We then combine these two saturation libraries. We transform the randomized DNA from the library into electrocompetent EBY-100 cells and grow the resulting $\sim 10^7$ transformants under selective conditions.

For the second generation, we begin by using the staggered extension process (STEP) (31) to recombine fastest 18 first-generation mutants. All 18 plasmids were mixed in equimolar amounts and then 1.25 ng of this mixture was used as a template for the STEP reaction with 200 mM dNTP, 150 nM each of the primers for the epPCR libraries described above, and 2.5 U of Taq polymerase (New England Biolabs). The reactions were cycled 99 times at 94 °C for 30 s followed by 55 °C for 5 s yielding a sharp band of the expected size at $\sim 100$ bp on an agarose gel. We used the product of the STEP reaction as a template for epPCR using the Mutazyme II kit following the standard protocol.

Preparation of Yeast for Sorting. HRP libraries were grown, induced, and purified as described above. For the second generation, we begin by using the staggered extension process (STEP) (31) to recombine fastest 18 first-generation mutants. All 18 plasmids were mixed in equimolar amounts and then 1.25 ng of this mixture was used as a template for the STEP reaction with 200 mM dNTP, 150 nM each of the primers for the epPCR libraries described above, and 2.5 U of Taq polymerase (New England Biolabs). The reactions were cycled 99 times at 94 °C for 30 s followed by 55 °C for 5 s yielding a sharp band of the expected size at $\sim 100$ bp on an agarose gel. We used the product of the STEP reaction as a template for epPCR using the Mutazyme II kit following the standard protocol.


